PCT

WORLD INTELLECTUAL PROPERTY ORGANIZATION International Bureau



INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT

(11) International Publication Number: WO 00/06706 C12N 9/00 (21) International Application Number: PCT/EP99/05413 (22) International Filing Date: 28 July 1999 (28.07.99) (23) International Filing Date: 28 July 1999 (28.07.99) (24) International Filing Date: 28 July 1999 (28.07.99) (25) International Filing Date: 28 July 1999 (28.07.99) (26) International Filing Date: 28 July 1999 (28.07.99) (27) International Filing Date: 28 July 1999 (28.07.99) (28) International Publication Number: WO 00/06706 (29) International Publication Number: 10 February 2000 (10.02.00) (21) International Publication Number: 10 February 2000 (10.02.00) (21) International Publication Number: 10 February 2000 (10.02.00) (22) International Publication Number: 10 February 2000 (10.02.00) (23) International Publication Number: 10 February 2000 (10.02.00) (24) Designated States: AE, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CU, CZ, DE, DK, EB, ES, FI, RB, GB, GR, EB, EB, EB, EB, EB, EB, EB, EB, EB, EB	INTERNATIONAL APPLICATION PUBLISH	HED U	UNI	DER THE PATENT COOPERAT	ION TREATY (PCT)
(21) International Application Number: PCT/EP99/05413 (22) International Filing Date: 28 July 1999 (28.07.99) Priority Data: 9816639.0 30 July 1998 (30.07.98) (71) Applicant (for all designated States except AT US): NOVARTIS AG [CH/CH]; Schwarzwaldallee 215, CH-4058 Basel (CH). PASZOWSKI, Jerzy [PL/CH]; Blauenweg 10, CH-4224 Nenzlingen (CH). (74) Agent: BECKER, Konrad; Novartis AG, Corporate Intellectual	(51) International Patent Classification 7:		(1	1) International Publication Number:	WO 00/06706
(22) International Filing Date: 28 July 1999 (28.07.99) (23) International Filing Date: 28 July 1999 (28.07.99) (24) International Filing Date: 28 July 1999 (28.07.99) (25) International Filing Date: 28 July 1999 (28.07.99) (26) International Filing Date: 28 July 1999 (28.07.99) (27) Priority Data: 9816639.0 30 July 1998 (30.07.98) (28) International Filing Date: 28 July 1999 (28.07.99) (29) International Filing Date: 28 July 1999 (28.07.99) (20) International Filing Date: 28 July 1999 (28.07.99) (20) International Filing Date: 28 July 1999 (28.07.99) (21) International Filing Date: 28 July 1999 (28.07.99) (22) International Filing Date: 28 July 1999 (28.07.99) (23) International Filing Date: 28 July 1999 (28.07.99) (24) International Filing Date: 28 July 1999 (28.07.99) (25) International Filing Date: 28 July 1999 (28.07.99) (26) International Filing Date: 28 July 1999 (28.07.99) (27) International Filing Date: 28 July 1999 (28.07.99) (28) International Filing Date: 28 July 1998 (30.07.98) (28) International Filing Date: 28 July 1998 (30.07.98) (29) International Filing Date: 29 July 1998 (30.07.98) (20) GE, GH, GM, HR, HU, ID, IL, IN, IS, IP, KE, KG, KP, KR, KZ, LC, LX, LR, LS, LT, LU, LV, MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, UA, UG, US, UZ, VN, YU, ZA, ZW, ARIPO patent (6H, GM, KE, LS, MW, SD, SL, SZ, UG, ZW), Eurasian patent (AT, AZ, BY, KG, KZ, MD, RU, TJ, TM), European patent (AT, BE, CH, CY, DE, DK, EE, KG, KR, KZ, LC, LK, LR, LR, LT, LU, LV, MD, MR, MN, MW, MX, NO, NZ, PL, PT, RO, RU, ST, EV, KG, KE, KG, KM, MN, MN, MW, MX, NO, NZ, PL, PT, RO, RU, ST, SK, SL, TJ, TM, TR, TT, UA, UG, US, VE, KG, KE, KG, KM, MN, MN, MW, MX, NO, NZ, PL, PT, RO, RU, ST, SK, SL, TJ, TM, TR, TT, UA, UG, US, VE, KG, KE, KG, KM, MN, MN, MN, MN, MN, MN, MN, MN, MN, M	C12N 9/00	A2	(4:	3) International Publication Date:	10 February 2000 (10.02.00)
Property, Patent & Trademark Dept., C11-4002 Basel (C11).	 (22) International Filing Date: 28 July 1999 (20) (30) Priority Data: 9816639.0 30 July 1998 (30.07.98) (71) Applicant (for all designated States except AT US): 1 TIS AG [CH/CH]; Schwarzwaldallee 215, CH-40 (CH). (71) Applicant (for AT only): NOVARTIS-ERFINDUNGE WALTUNGSGESELLSCHAFT MBH [AT/AT]; Strasse 59, A-1230 Vienna (AT). (72) Inventors; and (75) Inventors/Applicants (for US only): REVENKOVA, I [RU/CH]; Peter Rot-Strasse 76, CH-4058 Bas PASZOWSKI, Jerzy [PL/CH]; Blauenweg 10, Nenzlingen (CH). (74) Agent: BECKER, Konrad; Novartis AG, Corporate In 	28.07.9 G NOVAI 558 Bas EN VEI Brunn Ekaterin sel (CH CH 422	13 99) GB R-sel na H).	(81) Designated States: AE, AL, AM, BR, BY, CA, CH, CN, CU, CGD, GE, GH, GM, HR, HU, KP, KR, KZ, LC, LK, LR, LS, MN, MW, MX, NO, NZ, PL, SK, SL, TJ, TM, TR, TT, UA ZW, ARIPO patent (GH, GM UG, ZW), Eurasian patent (ARU, TJ, TM), European patent ES, FI, FR, GB, GR, IE, IT, I patent (BF, BJ, CF, CG, CI, CNE, SN, TD, TG). Published Without international search in	CZ, DE, DK, EE, ES, FI, GB, ID, IL, IN, IS, JP, KE, KG, LT, LU, LV, MD, MG, MK, PT, RO, RU, SD, SE, SG, SI, UG, US, UZ, VN, YU, ZA, KE, LS, MW, SD, SL, SZ, M, AZ, BY, KG, KZ, MD, (AT, BE, CH, CY, DE, DK, LU, MC, NL, PT, SE), OAPICM, GA, GN, GW, ML, MR,
	PASZOWSKI, Jerzy [PL/CH]; Blauenweg 10, (Nenzlingen (CH). (74) Agent: BECKER, Konrad; Novartis AG, Corporate In	CH-422	24 ial		

(54) Title: MAP KINASE PHOSPHATASE MUTANT

(57) Abstract

The present invention relates to DNA encoding proteins contributing to the regulation of a plant's response to DNA damage. DNA according to the invention comprises an open reading frame encoding a protein characterized by a stretch of amino acids or component amino acid sequence having 40% or more identity with an aligned component sequence of SEQ ID NO:3. Preferably the DNA encodes a MAP kinase phosphatase.

FOR THE PURPOSES OF INFORMATION ONLY

Codes used to identify States party to the PCT on the front pages of pamphlets publishing international applications under the PCT.

AL	Albania	ES	Spain	LS	Lesotho	SI	Slovenia
AM	Armenia	FI	Finland	LT	Lithuania	SK	Slovakia
AT	Austria	FR	France	LU	Luxembourg	SN	Senegal
ΑU	Australia	GA	Gabon	LV	Latvia	SZ	Swaziland
ΑZ	Azerbaijan	GB	United Kingdom	MC	Monaco	TD	Chad
BA	Bosnia and Herzegovina	GE	Georgia	MD	Republic of Moldova	TG	Togo
BB	Barbados	GH	Ghana	MG	Madagascar	TJ	Tajikistan
BE	Belgium	GN	Guinea	MK	The former Yugoslav	TM	Turkmenistan
BF	Burkina Faso	GR	Greece		Republic of Macedonia	TR	Turkey
BG	Bulgaria	HU	Hungary	ML	Mali	TT	Trinidad and Tobago
BJ	Benin	IE	Ireland	MN	Mongolia	UA	Ukraine
BR	Brazil	IL	Israel	MR	Mauritania	UG	Uganda
BY	Belarus	IS	Iceland	MW	Malawi	US	United States of America
CA	. Canada	IT	Italy	MX	Mexico	UZ	Uzbekistan
CF	Central African Republic	JP	Japan	NE	Niger	VN	Viet Nam
CG	Congo.	KE	Kenya	NL	Netherlands	YU	Yugoslavia
CH	Switzerland	KG	Kyrgyzstan	NO	Norway	zw	Zimbabwe
CI	Côte d'Ivoire	KP	Democratic People's	NZ	New Zealand	211	Zimbaowe
CM	Cameroon		Republic of Korea	PL	Poland		
CN	China	KR	Republic of Korea	PT	Portugal		
CU	Cuba	KZ	Kazakstan	RO	Romania		
CZ	Czech Republic	LC	Saint Lucia	RU	Russian Federation		
DE	Germany	Li	Liechtenstein	SD	Sudan		
DK	Denmark	LK	Sri Lanka	SE	Sweden		
EE	Estonia	LR	Liberia	SG	Singapore		

- 1 -

MAP kinase phosphatase mutant

The present invention relates to DNA encoding proteins contributing to the regulation of a plant's response to abiotic stress and in particular genotoxic stress.

Cells of all organisms have evolved a series of DNA repair pathways which counteract the deleterious effects of DNA damage and are triggered by intricate signal cascades. To be able to modify or improve DNA repair using gene technology it is necessary to identify key proteins involved in said pathways or cascades. Therefore it is the main object of the present invention to provide DNA comprising an open reading frame encoding such a key protein.

DNA according to the present invention comprises an open reading frame encoding a protein characterized by a stretch of amino acids or component amino acid sequence having 40% or more identity with an aligned component sequence of SEQ ID NO: 3. The protein characterized by SEQ ID NO: 3 is tracked down with the help of a T-DNA tagged Arabidopsis mutant showing hypersensitivity to methyl methanesulfonate (MMS). Said hypersensitivity as well as an observed hypersensitivity to other DNA damaging treatments such as UV light is indicative of the proteins' involvement in the repair of DNA damage, or in signaling pathways implicated in the response to similar genotoxic stress. The mutant is also sensitive to elevated temperature and anti-oxidant N-acetylcysteine. The mutant is not sensitive to osmotic shock, increased salinity, oxidative stress or elevated ehtylene levels. An important characteristic of the mutant is cell death in response to growth in small closed vessels. This phenotype can be complemented by addition of abscisic acid (ABA) to the growth media. Furthermore, the mutant is more sensitive to exogenously applied ABA compared with the wild type which supports the notion that the genes disclosed by the present invention (SEQ ID NO: 1) are involved in stress signaling mediated by ABA.

Sequence alignments of SEQ ID NO: 3 using commercially available computer programs such as BLASTP of the NCBI BLAST family of programs or TFASTA or BestFit of the Wisconsin Package Software, all based on well known algorithms for sequence identity or similarity searches, reveal that stretches of SEQ ID NO: 3 (component sequences) having more than 100 and preferably between 120 to 250 amino acids length can show between

- 2 -

20% and almost 40% sequence identity to aligned stretches of known phosphatases, particularly phosphotyrosine phosphatases, MAP kinase phosphatases or dual specificity phosphatases. Protein phosphatases are classified by their substrate specificities as either phosphoserine/threonine phosphatases (PSTPs) or phosphotyrosine phosphatases (PTPs). The dual specificity phosphatases (DSPs) dephosphorylate both phosphotyrosine and phosphoserine/threonine residues and represent a subfamily of PTPs. MAP kinase phosphatases (MKPs) belong to the family of DSPs. The sequence VHCCQGVSRS (SEQ ID NO: 4) found in SEQ ID NO: 3 can be interpreted as corresponding to the mammalian sequence motif IHCXAGXXRS (SEQ ID NO: 5) defining the family of PTPs, wherein the Ile at the first position can be replaced by Val and the Ser at the last position can be replaced by Thr

The present invention defines a new protein family the members of which are characterized by component amino acid sequences of more than 100 amino acid length showing 40% or higher amino acid sequence identity to aligned component sequences of SEQ ID NO: 3. Preferably said component sequences are of more than 120, more than 160 or even more than 200 amino acids length. The amino acid sequence identity is preferably higher than 50% or even higher than 55%. Most preferred are identities higher than 70%.

An example of DNA according to the present invention is described in SEQ ID NO: 1. The amino acid sequence of the protein encoded is identical to SEQ ID NO: 3. After alignment a stretch of the protein having about 140 amino acids shows 36% sequence identity to the MKP-1 protein described by Sun et al (Cell 75: 487-493, 1993). The identity determined after alignment with MKP-2 and MKP-3 is determined as 34% and 26%, respectively. Thus, according to the present invention a protein family related to MAP kinase phosphatases can be defined the members of which after alignment of a stretch of more than 100 amino acids length show 40% or higher amino acid sequence identity to SEQ ID NO: 3. Preferably, the amino acid sequence identity is higher than 50% or even higher than 55%. When making multiple sequence alignments, certain algorithms can take into account sequence similarities, such as same net charge or comparable hydrophobicity/hydrophilicity of the individual amino acids, in addition to sequence identities. The resulting values of sequence similarities, as compared to sequence identities, can help to assign a protein to the correct protein family in border-line cases. Proteins of particular interest, within the scope of the present invention, are MAP kinase phosphatases the amino acid s quence of which comprises at least one of the following characteristic amino acid subsequences:

. . . .

G - 3 -

(a) TSILYDVFDYFEDV (SEQ ID NO: 6)
(b) FVHCCQGVSRST (SEQ ID NO: 7)
(c) FVHC (SEQ ID NO: 8)
(d) QGVSRS (SEQ ID NO: 9)
(e) YFKSD (SEQ ID NO: 10)

DNA encoding proteins belonging to the new protein family according to the present invention can be isolated from monocotyledonous and dicotyledonous plants. Preferred sources are corn, sugar beet, sunflower, winter oilseed rape, soybean, cotton, wheat, rice, potato, broccoli, cauliflower, cabbage, cucumber, sweet corn, daikon, garden beans, lettuce, melon, pepper, squash, tomato, or watermelon. However, they can also be isolated from mammalian sources such as mouse or human tissues. The following general method, can be used, which the person skilled in the art will normally adapt to his specific task. A single stranded fragment of SEQ ID NO: 1 or SEQ ID NO: 2 consisting of at least 15, preferably 20 to 30 or even more than 100 consecutive nucleotides is used as a probe to screen a DNA library for clones hybridizing to said fragment. The factors to be observed for hybridization are described in Sambrook et al, Molecular cloning: A laboratory manual, Cold Spring Harbor Laboratory Press, chapters 9.47-9.57 and 11.45-11.49, 1989. Hybridizing clones are sequenced and DNA of clones comprising a complete coding region encoding a protein with more than 40% sequence identity to SEQ ID NO: 3 is purified. Said DNA can then be further processed by a number of routine recombinant DNA techniques such as restriction enzyme digestion, ligation, or polymerase chain reaction analysis.

The disclosure of SEQ ID NO: 1 enables a person skilled in the art to design oligonucleotides for polymerase chain reactions which attempt to amplify DNA fragments from templates comprising a sequence of nucleotides characterized by any continuous sequence of 15 and preferably 20 to 30 or more basepairs in SEQ ID NO: 1. Said nucleotides comprise a sequence of nucleotides which represents 15 and preferably 20 to 30 or more basepairs of SEQ ID NO: 1. Polymerase chain reactions performed using at least one such oligonucleotide and their amplification products constitute another embodiment of the present invention.

Knowing the nucleotide sequence of the Arabidopsis MKP1 gene and the amino acid sequence of the encoded protein it is possible to identify proteins interacting with AtMKP1 and to clone their corresponding genes using well known techniques. For example radioactively labeled AtMKP1 protein can be used for interactive cloning on cDNA expression libraries. AtMKP1 protein or parts thereof can be used to generate polyclonal or monoclonal antibodies specific for AtMKP1. The AtMKP1 gene can be used to generate variants of AtMKP1 protein tagged with GST, MYK or His. Said antibodies and MKP1 variants allow to isolate native protein complexes by immunoprecipitation and to determine sequences of proteins present in these complexes by micro-sequencing. The resulting sequence information can in turn be used to clone corresponding genes. Alternatively, said antibodies or tagged MKP1 variants can be used to screen epitope libraries for epitopes which interact with AtMKP1 protein. The AtMKP1 protein and parts thereof, in particular the N-terminal 490 amino acid region and the C-terminal 492 amino acid region can also be used to search for interacting proteins with a Two-hybrid system (e.g. in yeast, in mammalian cells, or in bacteria). This allows to obtain sequence information about interacting proteins.

Based on the disclosed finding that AtMKP1 proteins are involved in a plant's abiotic environmental stress response, it becomes possible to engineer the corresponding signaling pathway, of which AtMKP1 is a part, to be chemically regulated due to chemical activation or repression of transgenes encoding AtMKP1 or proteins interacting therewith. Such plants can be obtained by transformation with the corresponding genes under control of chemically inducible promoters. Application of inducers is expected to modify the activity of the AtMKP1 signaling pathway and to result in altered adaptation to abiotic environmental stress. Alternatively, AtMKP1 protein or its interacting proteins can be used as targets for chemicals inhibiting or stimulating their activities which again is expected to modify abiotic stress responses.

EXAMPLES:

Example 1: Cloning of the gene responsible for the mkp1 mutant phenotype

Arabidopsis T-DNA insertion lines as produced by the INRA-Versailles and available from the Nottingham Arabidopsis Stock Center (NASC) are screened for sensitivity to methyl methanesulfonate (MMS) at a concentration of 100 ppm as described by Masson et al,

· · - 5 -

Genetics 146: 401-407, 1997. Plants which die in the presence of 100 ppm MMS are found in the family AAN4. Thus, the corresponding T-DNA insertion mutation is assumed to give rise to this hypersensitive phenotype. This assumption is supported by genetic analysis showing co-segregation of the hypersensitive phenotype with the T-DNA insertion.

Genomic DNA from the mutant plants is isolated as described by Dellaporta et al, Plant Mol Biol Reporter 1: 19-21, 1983. A fragment of genomic DNA flanking the right border of the inserted T-DNA is rescued essentially according to Bouchez et al. Plant Mol Biol Reporter 14: 115-123, 1996, with minor modifications. Genomic DNA is digested with Pstl, ethanol precipitated and resuspended in H₂O. DNA of vector pResc38 (Bouchez et al supra) is digested with Pstl and dephosphorylated with shrimp alkaline phosphatase. The phosphatase is heat inactivated, the vector DNA is ethanol precipitated and resuspended in H₂O. 2.5 μg of PstI digested genomic DNA and 2.5 μg of PstI digested and dephosphorylated vector are mixed and ligated overnight at room temperature in a total volume 100 µl in the presence of 10 units of T4 DNA ligase. The DNA of the ligation mixture is precipitated with ethanol, resuspended in 50 µl H₂O, and digested with Xbal in a total volume of 100 μl. Xbal digested DNA is precipitated with ethanol and resuspended in H₂O. A second overnight ligation reaction in the presence of 10 units T4 DNA ligase is performed in a total volume of 200 µl at room temperature to achieve circularization of DNA fragments. The DNA of the ligation mixture is again precipitated with ethanol, rinsed two times with 70% ethanol, dried and dissolved in 5 μl H₂O. Two 2 μl aliquots are used for electroporation of electrocompetent E.coli XL1-Blue cells (Stratagene) according to the manufacturer's instructions. Transformants containing the inserted T-DNA and adjacent Arabidopsis genomic DNA sequences are selected on plates with 50mg/l kanamycin. Single bacterial colonies are analyzed by isolation of plasmid DNA using QIAprep Spin Plasmid Kit (Qiagen) and restriction digestion with Pstl and Xbal. Plasmid pBN1 containing 3.7 kb of inserted T-DNA linked to 5 kb of Arabidopsis DNA is identified. Sequencing of the junction site is performed using a primer directed towards the flanking plant DNA and having the nucleotide sequence 5'-GGTTTCTACAGGACGTAACAT-3' (SEQ ID NO: 14) complementary to T-DNA 41 nucleotides from the right border. Digestion of this clone with Sstl allows isolation of a 960 bp fragment which when labelled with 32P can be conveniently used as a probe to screen wild type Arabidopsis genomic and cDNA libraries in order to identify the wild type gene affected in the mkp1 mutant line.

Example 2: Cloning of the AtMKP1 wild-type gene

The 960 bp Sstl fragment mentioned at the end of example 1 is labeled with ³²P by random oligonucleotide-primed synthesis (Feinberg et al, Anal Biochem 132: 6-13, 1983) for use as a probe in the following hybridization experiments.

Southern blot analysis of *Arabidopsis* wild type and *mkp1* DNA digested with EcoRV confirms that in the *mkp1* genomic DNA the sequence hybridizing to the probe is linked to T-DNA.

Northern blot analysis of *Arabidopsis* wild type RNA reveals the presence of a hybridizing transcript in RNA extracted from seven-day-old wild type seedlings. No such hybridizing fragment is detected in the corresponding RNA of *mkp1* seedlings.

A <u>cDNA library</u> (Elledge et al, 1991) and a <u>genomic library</u> (Stratagene) of wild type *Arabidopsis thaliana* ecotype Columbia is screened with the labelled SstI fragment mentioned above. Screening of the bacteriophage λ libraries is performed according to the protocols described in chapter 6 of Ausubel et al, 1994, "Current protocols in molecular biology", John Wiley & Sons, Inc. Hybridization is performed as described by Church and Gilbert, Proc Natl Acad Sci USA 81: 1991-1995, 1984. Bacteriophage clones hybridizing to SstI fragment are subjected to *in vivo* excision of plasmids according to Elledge et al, Proc Natl Acad Sci USA 88: 1731-1735, 1991, and Stratagene protocols. Inserts of the plasmids obtained are further analyzed by sequencing.

By partial sequencing and alignment of ten overlapping clones (pBN5.1 to pBN5.10) isolated from the genomic library a continuous genomic sequence of 6356 bp (see SEQ ID NO: 1) is decoded.

Ten cDNA clones representing the same gene, one of them a 3.0 kb full-length cDNA (SEQ ID NO: 2), are isolated from the cDNA library.

Example 3: Sequence Analysis and Alignments

The 3 kb full-length cDNA clone of SEQ ID NO: 2 encodes an ORF with the start codon being defined by basepairs 298-300 and the stop codon by basepairs 2650-2652. The ORF encodes a protein consisting of 784 amino acids (SEQ ID NO: 3) and a predicted molecular mass of 86.0 kD. Alignment with the genomic sequence of SEQ ID NO: 1 reveals three introns. T-DNA is inserted within the coding sequence of the *mkp1* mutant DNA before basepair position 502 according to the numbering of SEQ ID NO: 2. The sub-sequence

WO 00/06706 PCT/EP99/05413

· - 7 -

VHCCQGVSRS (SEQ ID NO: 4) found in SEQ ID NO: 3 can be interpreted as corresponding to the mammalian sequence motif IHCXAGXXRS (SEQ ID NO: 5) defining the family of protein tyrosine phosphatases, wherein the IIe at the first position can be replaced by Val and the Ser at the last position can be replaced by Thr (Van Vactor et al, Curr Opin Gen Dev 8: 112-126, 1998). Therefore it is concluded that the wild type ORF encodes a protein tyrosine phosphatase with invariant aspartic acid, cysteine, and arginine residues (Fauman et al, Trends Biochem Sci 21: 413-417, 1996) in positions 204, 235 and 241 according to SEQ ID NO: 3.

A data base search using the TFASTA program (Wisconsin Package Version 9.1, Genetics Computer Group (GCG), Madison, Wisc.) reveals that the encoded phosphatase has a significant similarity to dual specificity phosphatases. The closest homologue identified is *Xenopus laevis* MAP kinase phosphatase (MKP; Lewis et al, J Cell Sci 108: 2885-2896, 1995) showing 38.1% identity and 52.5% similarity in a 140 amino acid overlap region. The deduced AtMKP1 protein also has 36.0% identity and 52.5% similarity with a 140 amino acid overlap region encoded by the rat 3CH134/CL100 cDNA representing a rat MKP1. Essentially identical results are obtained when using the BLASTP 2.0.4 (Feb-24-1998) program of the NCBI BLAST family of programs which, allowing gapped alignment, compares an amino acid query sequence against a protein sequence database (Altschul et al, Nucleic Acids Res. 25: 3389-3402, 1997). No higher plant homologues are identified. The genomic position of the *AtMKP1* gene is determined by hybridization to filters containing genomic YAC clones publicly available from the Arabidopsis Biological Resource Center (Ohio, USA). The *AtMKP1* gene is found to map to chromosome 3 between markers ve022 and BGL1.

Example 4: Complementation

mkp1 mutant plants are transformed with DNA comprising the corresponding wild type genomic DNA including promoter and polyadenylation signal to find out whether the cloned wild type gene is able to complement the mutant mkp1 phenotype.

mkp1 mutant plants harbor T-DNA containing the NPTII and bar marker genes under the control of nos and CaMV35S promoters, respectively. Therefore, different marker genes are used for the transformation construct. The vector used is a derivative of p1'barbi which is

highly efficient in Arabidopsis transformation (Mengiste et al, Plant J 12: 945-948, 1997). In

p1'barbi the EcoRl fragment containing 1'promoter, bar gene coding region, and CaMV 35S polyadenylation signal is inverted in respect to the T-DNA borders by EcoRI digestion and re-ligation. In the resulting plasmid the 1'promoter (Velten et al, EMBO J 3: 2723-2730. 1984) is directed towards the right border of the T-DNA. This plasmid is digested with BamHI and Nhel, and the bar gene and CaMV 35S polyadenylation signal are replaced by a synthetic polylinker with the sites for the restriction enzymes BamHI, HpaI, ClaI, StuI and Nhel. The resulting plasmid is digested with BamHI and Hpal and ligated to a BamHI-Pvull fragment of pROB1 (Bilang et al, Gene 100: 247-250, 1991) containing the hygromycin-Bresistance gene hph linked to the CaMV 35S polyadenylation signal. The T-DNA of the resulting binary vector p1'hygi contains the hygromycin resistance selectable marker gene under the control of the 1'promoter and unique cloning sites for the restriction enzymes Clai, Stul and Nhel located between the marker gene and the T-DNA right border. p1'hygi is used to insert the reconstructed AtMKP1 gene as follows. Plasmid pBN1 of example 1 is digested with Pstl and Munl and dephosphorylated. The restriction fragment containing the 3'portion of the AtMKP1 gene and pBluescript-SK(+) is purified from the agarose gel and ligated to the Pstl-Muni restriction fragment of the wild type genomic clone pBN5.2 (example 2) including the 5' end of the coding sequence of the AtMKP1 gene and 2.4 kb of upstream sequences. The reconstructed AtMKP1 gene is excised by Pstl and Notl and after filling the ends is inserted into the Stul site of p1'hygi. The construct is introduced by transformation into Agrobacterium tumefaciens strain C58CIRif^R containing the nononcogenic Ti plasmid pGV3101 (Van Larebeke et al. Nature 252: 169-170, 1974), T-DNA containing the reconstructed AtMKP1 gene is transferred to mutant plants by the method of in planta Agrobacterium mediated gene transfer (Bechtold et al, C R Acad Sci Paris, Life Sci 316: 1194-1199, 1993). Seeds of infiltrated plants are grown on hygromycin-containing medium to screen for transformants. The progeny of selfed hygromycin resistant plants is analyzed for the segregation of hygromycin resistance. The families in which a 3:1 segregation ratio is observed are used to isolate homozygous lines bearing the newly introduced T-DNA inserted at a single genetic locus. The obtained hygromycin resistant lines are analyzed by Northern blot analysis for the restoration of AtMKP1 expression. In these lines the restoration of transcription of the AtMKP1 gene can be observed as well as the restoration of the wild typ level of MMS resistance and ABA mediated stress responses. Complementation is not observed in plants transformed with p1'hygi only.

9 - 9 -

Example 5: Cloning of homologous sequences from other plant species

Use of AtMKP1 cDNA as a probe for Southern hybridization with genomic DNA from other plant species such as Sinapis alba (mustard), Lycopersicum esculentum (tomato) and Zea mays (maize) is successful in the case of Sinapis alba which belongs to the same family as Arabidopsis (Brassicaceae).

Homologous sequences from the other species can be identified in a PCR approach using degenerate primers 1-3 below, wherein I is inosine, derived from the regions conserved between VH-PTP13 of *Clamydomonas eugametos* and AtMKP1 protein:

Primer 1 (forward): 5'-AAY AAY GGI ATH ACI CAY ATH YT-3' (SEQ ID NO: 11); Primer 2 (reverse): 5'-YTG RCA IGC RAA ICC CAT RTT IGG-3' (SEQ ID NO: 12); Primer 3 (reverse): 5'-IGT CCA CAT IAR RTA IGC DAT IAC (SEQ ID NO: 13);

A PCR reaction is performed in a total volume of 50 μ l containing 1x reaction buffer (Qiagen), 200 μ M of each dNTP, 1.25 units of Taq polymerase (Qiagen), and 100 pmol of each primer.

Reaction 1 is performed with primers 1 and 2 using genomic DNA from *Sinapis alba* (200 ng), *Lycopersicum esculentum* (400 ng), or *Zea mays* (600 ng) as the original template DNA. Amplification is carried out after an initial denaturation step of 3 min at 94°C, followed by 30 cycles of 30 sec at 94°C, 30 sec at 40°C, and 3 min at 72°C. The resulting amplification mixture is diluted 10³ fold.

Reaction 2 is performed using 2µl of the above dilution to provide the necessary template DNA. This time primers 1 and 3 are used under the same conditions as specified for reaction 1. The resulting amplification products are cloned into the T/A vector pCR2.1 (Invitrogen) and further analyzed by nucleotide sequencing.

Using this PCR approach it is possible to amplify sequences homologous to the *AtMKP1* gene from all the species mentioned above. Whereas the nucleotide sequence from *Sinapis alba* SaMKP1 ((SEQ ID NO: 15 encoding SEQ ID NO: 16) is 90.8% identical to the *AtMKP1* sequence, the nucleotide sequence from *Lycopersicum esculentum LeMKP1* (SEQ ID NO: 17 encoding SEQ ID NO: 18) is 72.3% and the *Zea mays* sequence

ZmMKP1 (SEQ ID NO: 19 encoding SEQ ID NO: 20) 71.8% identical. The fragments hybridize to genomic DNA from corresponding species under the usual hybridization conditions for Southern blot analysis. The fragments can be used as probes to screen cDNA libraries for corresponding cDNA sequences.

The 243 bp ZmMKP1 fragment amplifying from maize DNA is used as a probe to screen a maize cDNA library (Clontech) made in the Lambda ZAP®II Vector (Clontech) from "Blizzard" hybrid etiolated shoots, which were treated with the herbicide safener Benoxacor. The titer of the library is determined as 3×10⁹ pfu/ml.

Library screening is conducted as described in the Clontech Lambda Library Protocol Handbook, with some slight modifications. Briefly, a single colony of XL-1 Blue is picked and incubated overnight at 37°C in LB medium, containing 10mM MgSO₄ and 0.2% maltose. 600 µl of stationary phase grown bacteria for each 150 mm plate is combined with 100 μl of phage library dilution in sterile 1× lambda dilution buffer (100mM NaCl; 10mM MgSO₄; 35mM Tris-HCl, pH7.5) to yield approximately 30,000 pfu per plate. This mixture is incubated at 37°C for 15 minutes, subsequently 7 ml of melted LB soft top agarose (at 48°C) is added to the cell suspension for each 150 mm plate, shortly mixed and then poured on two-day-old LB^{MgSO4} agar plates, which have been pre-warmed to 37°C for four hours. The plates are then incubated at 37°C until plaques reach appropriate sizes (after about 8 to 9 hours). After chilling the plates at 4°C for one hour, phage particles are transferred to Hybond N nitrocellulose filters and the orientation of each filter to its plate is recorded with a waterproof pen. The filters are then treated by placing them on Whatman 3MM paper saturated with the appropriate solution. The treatments include denaturation solution (0.5M NaOH; 1.5M NaCl) for 2 minutes, followed by neutralization solution (0.5M Tris-HCl, pH 7.2; 1.5M NaCl; 1mM EDTA) for 3 minutes and 2× SSC for 3 minutes. DNA is subsequently crosslinked to the filters by UV.

Filters are then pre-hybridized, hybridized with the radioactivelly labeled *ZmMKP1* fragment and washed as described in Sambrook et al, Molecular cloning: A laboratory manual, Cold Spring Harbor Laboratory Press, chapters 9.47-9.57 and 11.45-11.49, 1989. An agar plug from the position of a positive plaque is then removed from the master plate and incubated overnight at 4°C in 1 ml of 1× lambda dilution buffer, containing 20 µl of chloroform. Each titer is determined and the phages are re-plated to obtain approximately 200 to 500 plaques on a 150 mm plate for a secondary screen as described above. Single plaques of interest

WO 00/06706 PCT/EP99/05413

, 🤧 . - **11** -

are collected from the agar plates and incubated over night at 4°C in 500 μ l of 1× lambda dilution buffer and 20 μ l of chloroform.

The pBluescript phagemid is excised from the λZAP™ vector as described by the *In Vivo* Excision Protocol using the ExAssist/SOLR System in the Stratagene Uni-ZAPTM XR Library Instruction Manual (1993). A 1/100 dilution is made of XL1-Blue MRF' and SOLR overnight cultures (at 30°C) and incubated at 37°C for 2-3 hours. XL1-Blue MRF' cells are then pelleted for 10 minutes at 1,500×g and re-suspended at an OD₆₀₀ = 1.0 in 10mM MgSO₄. 200 ul of these XL1-Blue cells are then combined with 250 ul of phage stock and 1 ul of ExAssist helper phage in a 50 ml conical tube and incubated at 37°C for 15 minutes. 3 ml of LB broth is added and incubated at 37°C for 5 hours, after which the cells are pelleted for 15 minutes at 2,000×g and the supernatant transferred to a new tube. The tube is then heated at 70°C for 15 minutes and centrifuged again for 15 minutes at 4,000×g. The supernatant, containing the excised phagemid pBluescript packaged as filamentous phage particles, is decanted into a new tube. 10 and 100 µl of this phage stock are then added to two tubes with 200 μ i of SOLR cells that have been allowed to grow to OD₆₀₀ = 0.5-1.0 before being removed from the incubator and further incubated at room temperature. The tubes are incubated at 37°C for 15 minutes, followed by plating 10-50 µl from each tube on LB^{amp} (50 μg/ml) and over night incubation at 37°C.

The positive clones are checked for insert size by *EcoRI/Xho*I double digestion and end-sequencing with T3 and T7 promoter primers (Promega).

Screening of 360,000 pfu of the library results in three identical clones of 2.2 kb containing the 3' poly(A) tail, but lacking part of the 5' end, including the translation initiation site. The gene corresponding to the identified partial cDNA clone is named ZmMKP2, as it is not identical with the ZmMKP1 fragment used as the probe (92.3% identity on the nucleotide level over the 196 bp fragment flanked by the primers 1 and 3). An additional 213 nucleotides are amplified and cloned by 5' RACE (rapid amplification of cDNA ends) carried out following the instructions of the 5'/3' RACE Kit (Boehringer Mannheim) resulting in a longer cDNA sequence of 2,452 bp but still not complete, judged by the predicted mRNA length from the RNA gel blot analysis and the absence of a possible translation initiation site.

The sequence information gained from the *ZmMKP2* cDNA including the additional 213 nucleotides obtained by 5' RACE (SEQ ID NO: 21 encoding SEQ ID NO: 22) is used to

design two additional backward oriented degenerate primers wherein I is inosine to 3' regions conserved between the deduced peptide sequences of *ZmMKP2* and *AtMKP1*:

```
Primer 4 (reverse): 5'-GCI GCY TTI GCR TCY TTY TCC-3' (SEQ ID NO: 25);

Primer 5 (reverse): 5'-YTC ICK IGC IGG IAR RTG IGT YTC-3' (SEQ ID NO: 26)
```

These primers are used to PCR amplify a larger fragment of a MAP kinase phosphatase gene from tomato. The amplified and cloned 522 bp long fragment is not identical to *LeMKP1*. Therefore, its corresponding gene is named *LeMKP2* (SEQ ID NO: 23 encoding SEQ ID NO: 24; 75% identity on the nucleotide level over the stretch of 196 bp of *ZmMKP1* flanked by primers 1 and 3). The origins of all identified MAP kinase phosphatase homologous gene sequences are confirmed by Southern blot analysis.

The following Table shows an alignment of a continuous stretch of 312 amino acids of *AtMKP1* with the related amino acid sequence of *ZmMKP2*.

AtMKP1	139	KREKIAFFDKECSKVADHIYVGGDAVAKDKSILKNINGITHILNCVGFICP 188
ZmMKP2	24	:::.
AtMKP1	189	EYFKSDFCYRSLWLQDSPSEDITSILYDVFDYFEDVREQSGRIFVHCCQG 238
ZmMKP2	74	EYFKSDLVYRTLWLQDSPTEDITSILYDVFDYFEDVREQGGRVLVHCCQG 123
AtMKP1	239	VSRSTSLVIAYLMWREGQSFDDAFQYVKSARGIADPNMGFACQLLQCQKR 288
ZmMKP2	124	
AtMKP1	289	VHAFPLSPTSLLRMYKMSPHSPYDPLHLVPKLLNDPCPGSLDSRGAFIIQ 338
ZmMKP2	174	- : : : VHAIPLSPNSVLRMYRMAPHSQYAPLHLVPKMINDPSPATIDSRGAFIVH 223
AtMKP1	339	LPSATYTWVGRQCETTMEKDAKAAVCQTARYEKVEAPTMVVRBCDEPVYY 388
ZmMKP2	224	VLSSLYWWGMKCDPVMEKDAKAAAFQVVRYEKVQGHIKVVREGLEPQEF 273
AtMKP1	389	WDAFASILPMIGGSVIKVQPGDRKVDAYNLDFEIFQKAIE 428
ZmMKP2	274	
AtMKP1	429	GGFVPTLASSNNEHETHLPARE 450
ZmMKP2	324	GGVVPAFSTSGAGDETHLPARE 345

What is claimed is:

- A DNA comprising an open reading frame encoding a protein characterized by a component amino acid sequence having 40% or more identity with an aligned component sequence of SEQ ID NO: 3
- 2. The DNA according to claim 1 comprising an open reading frame encoding a plant MAP kinase phosphatase.
- The DNA according to claim 1 wherein the open reading frame encodes an amino acid sequence selected from the group of amino acid sequences described in SEQ ID NO:
 SEQ ID NO: 7, SEQ ID NO: 8, SEQ ID NO: 9, and SEQ ID NO: 10.
- 4. The DNA according to claim 1, wherein the open reading frame encodes a protein characterized by the amino acid sequence of SEQ ID NO: 3
- The DNA according to claim 1 characterized by the nucleotide sequence of SEQ ID NO: 1
- 6. The DNA according to claim 1 wherein the open reading frame encodes a protein contributing to repair of DNA damage in a plant cell.
- 7. The DNA according to claim 1 wherein the open reading frame encodes a protein conferring hypersensitivity to treatment with methyl methanesulfonate (MMS).
- 8. The DNA according to claim 7 wherein the open reading frame encodes a protein conferring hypersensitivity to treatment with UV light or X-rays.
- 9. The DNA according to claim 7 wherein the open reading frame encodes a protein interfering with abscisic acid signal transduction.
- 10. The protein encoded by the open reading frame of any one of claims 1 to 9.
- 11. A method of producing DNA according to claim 1, comprising
 - screening a DNA library for clones which are capable of hybridizing to a fragment of the DNA defined by SEQ ID NO: 1, wherein said fragment has a length of at least 15 nucleotides;
 - sequencing hybridizing clones;
 - purifying vector DNA of clones comprising an open reading frame encoding a protein with more than 40% sequence identity to SEQ ID NO: 3
 - optionally further processing the purified DNA.
- 12. A polymerase chain reaction wherein at least one oligonucleotide used comprises a sequence of nucleotides which represents 15 or more basepairs of SEQ ID NO: 1.

- 1

SEQUENCE LISTING

```
<110> Novartis AG
<120> Organic compounds
<130> Silencing Gene
<140>
<141>
<160> 26
<170> PatentIn Ver. 2.1
<210> 1
<211> 6356
<212> DNA
<213> Arabidopsis thaliana
<400> 1
gttcaaggtg gtgttatttn cagtattaga aaagaggctt ctagagagag tttctaaatt 60
atattttgaa cacggccgga ttggttgtct tatgattaag tgaatgcttt agatctggtg 120
acgattggta tatgagatat atagtanaat gatagaacat cagaaatact ataagtcacc 180
atatttttaa aaaataaatg ctatagaatt tttgttttgg taattgttat aactctacaa 240
agttatgtgt tatagagttt tactagtttc atcgttatca tgtgtatagt ataacncaac 300
aaaagaaatt ttaaatatct gaaaacataa aaatttaata aaatgatgtg agtataagaa 360
aaaaagaaag aaagaaacaa cgtaaaaaat aaaaatcatt catacatata acaattttca 420
aaagatcaat gttaacttta attagtettt tttettatgt tteatgeaaa tgeaatagta 480
ttacttttct ttaatctaag attatgtgtt gcttttaagc aagaactatt caaagagtca 540
aagacatgca tgtaaacttt agagaacggg atctttcatc aatcttattt ttttcccttt 600
tttttttaca acgaacgata ggtagtaagc atgaatctgg ttcaaagttt ggatcagtgt 660
aageteatag tietgigtag aaaaaaegig gagatiggag agaaattaet aagagagega 780
agagaagagt aaacctagaa gatacaaaag acgctatcaa aagattgttt tatgtgttta 840
tgaaacattc acttaggcca gtataaatat taaatgtcat tttgagaaag aaaaaaaaat 900
ggtgtcattt tgaggatata aatattttct caacaagatg ttttagactt tcaacataac 960
gatcatttaa aactacataa tttatctctt cgttaaaaac ttaacattaa caaaaagtta 1020
aattcgacaa gaagagttta ttcaataagt atactgtatg tgtttatttt attcacatga 1080
aaaqtotcaa acaacacttg ggatacatat ttcatgccta aaaatcgtat aacaacattt 1140
aaggtaatca gtaaatttga tgcctaaact aaaatatgtg accattttga ccaatgagca 1200
ttttttggaa ataaaactac tgcatacgac ctggtcaaac acatgaactg ccactgcttt 1260
taaaactacc tttttaaact tttgacgtcc atttttattt actaattatt atgttaatca 1320
aaccactaac attatgatte gaaatttega gtgatgattq ttagaaqtga tqtgaatgat 1380
geggtataaa acaaattgtg atatatattc actcatatat atcaaaatca aaatagetta 1440
tegttecaaa acatgattga taaaatgtaa ttactattea aaaettaaag ceagtagtta 1500
atgaggaaaa gtctttttt ttttttcttt ttttttctca tagttaaaag acataaaaaa 1620
aattaatatt acaaacaaga caaaaaaaaa gaataaataa aacaacaatc attactgtca 1680
aatcaattat cagagggaaa agttattaag aaatgtcaac caatgagtat cattatcatc 1740
atgetttttt agaccettet tittaattea teaagatita gtettgitta taaattigaa 1800
gtaaaattot aaatgaatca attotacaat tttttcccta cattattgta acgataaaat 1860
ttaagatcaa caattacttc gttaattttt tttctgataa aaatatttga tatctttctg 1920
attatgatat attagcattg tttttgtatt gtatgtatgg taattaattt tagttcaaaa 1980
```

gaataataaa tggtttgatt agcatgaatt taataaaaaa ataagactga ataaacatag 2040 gtaataaact ttgtttcttt tggtaaatgt aaaattagaa aaaatcataa tcccaaggta 2100 attactataa ttacattcaa tgtcagaatt aaacgtagtg aataaaaacc gtaaaacatg 2160 agaaaaacaa gaatttatta tetttgacaa geaataaatg aaatgetgac aaaaaattgg 2220 tttcaaagtt tcaacgcgtt tcttataata agaattcaat ttcgtgcagc taatcaggag 2280 ataattatca taattaaatt aatcgttact actttataat actcccaaaa taatcgaaaa 2340 cgaatttatt ttattgtaat ttgtttaata ataaagaatt actgtttcct cccacttccc 2400 atototottt toottitigtg tiottottot totoogottt gtitooccaa totototot 2460 ctctctctct gaagaaaat aaataaaaga tctaactttg acggctctct taatcttact 2520 cacteegtaa gtteeaaate teteteettt aeteteatat etatategte egaacaaaac 2580 ccaggagaat tgcttcaccc cctttttggg tttttaatca ttttctcaga ttctcagttt 2640 ctgtttccgt cttctagatt ctgggttcag tttctgtttt gctcttattg aattttctta 2700 ttcattttgt gtttcggagt tattcatggt agctgaattt gttaattctg atgttgtttt 2760 gegttttett ettttetagt ttggetatgt egtetttgat etgatgetgg gttattetet 2820 ttccctctgt tttggtttct tttagggttt taagtcggaa tagactgatg ggagcttgat 2880 ggttattgtt agatcagatg tggatttaaa gccttcgctg aactaacaag tctatggaag 2940 aagcaaagac cettgtttta cactgtatgt tgtgaggaat ttgtctgatt ttgggtgata 3000 aaggtgaagt ctttgagttt gtaattttga gataagattg gatggtggga agagaggatg 3060 cgatggggaa tgatgaagct cctcctggtt ctaagaaaat gttttggcgg tctgcctctt 3120 ggtctgcttc acggactgca tcacaagttc ctgagggtga tgagcaaagc ctgaacattc 3180 cgtgtgctat tagttctggg ccgagtcgaa gatgtccagc tgctcctttg acacctcgtt 3240 cacatcataa cagcaagget agagettgtt tgccaccatt gcagcetett gccatttcta 3300 ggaggagett agacgagtgg cetaaggegg gtteggatga tgteggtgag tggeeteate 3360. caccaacacc tagegggaac aaaacegggg agagattgaa getegattta teateaacge 3420 agcagcgggt aacagataag agctctggtc tagctaagag ggagaagatt gctttctttg 3480 ataaagaatg ttcaaaggtt gctgatcata tatatgtggg tggagatgct gtggcgaaag 3540 acaagagcat actgaaaaac aatggaatca cgcatatctt gaactgcgtt ggttttatct 3600 gtccggaata tttcaagtct gatttttgtt acagatcctt gtggttacag gatagtccgt 3660 cagaggatat agctagtatt ctgtacgatg tgtttgacta ctttgaagat gtgagggagc 3720 aaagtggaag gatctttgtt cattgttgtc aaggggtttc acgatctacc tcgttggtaa 3780 tagcatatct gatgtggaga gaagggcaaa gttttgatga tgcatttcag tatgtgaagt 3840 ctgctagagg tattgctgat cctaacatgg gctttgcttg ccaattgtta caatgccaaa 3900 agagggttca tgcgttcccg cttagcccta cctccttact tagaatgtac aaaatgtctc 3960 cacactetee tratgaceer tracatetry treeaaaact grigaargar ceargeeega 4020 gttctctgga ttcaagaggt gcatttatca ttcagttacc ttctgcaatt tacatttggg 4080 ttggtaggca gtgtgaaacc atcatggaga aagatgcaaa agctgctgtt tgtcagattg 4140 ctcgatatga gaaagtcgaa gcacctatta tggtggtcag agaaggtgat gagcctgttt 4200 attactggga tgcatttgca agcattttgc ctatgattgg gggctcggta attaaagttc 4260 aaccaggtga caggaaggtc gacgcatata atctggattt tgagattttt cagaaagcca 4320 tagagggagg ttttgtgcca actttagcat catccaacaa cgaacatgag actcatcttc 4380 ctgcaaggga aaacagttgg agctcactga aatgtaagtt tgcatcaagg tttgacaaag 4440 gttttcggta tgtctccaaa acgccactat ctagggtcta ttcagattcg atgatgatcg 4500 / tgcattcatc aggeteacet tecteaacaa ettetteate atecaetgeg tegeeteett 4560 ttetetetee egattetgta tgtteaacaa atteaggeaa tagettaaag agtttetete 4620 aatcototgg acgttcgtcc ttgagacett ctattccacc atcgctgaca ttgcctaaat 4680 tttccagect ateceteete cetteecaaa etteteetaa agaatetegt ggtgtcaata 4740. ettttettea acegteacea aatagaaagg etteacette tettgetgag egtagaggea 4800 gcctgaaagg atctctgaag ttgccaggtt tggctgattc caacagaggc acacctgctt 4860 ttactttaca tccggatgat agtaatgaca tagtcttcaa tctggagggt attagaaacg 4920 gcgatctata tccaccaagt gattgcaaag ggacaagtgt agattcagat ttgccagaga 4980 aggaaattat atccttaatc agttgcagta aatctgacag acacaaatcg ggaggtgata 5040. ctgataget: tggccageet ttagcatgte gttggccaag tatggagatg attacaaaac 5100 tgagcagagc ttacttagat tcagaatctg ttatagcaat cccgttgcca agtgatgctg 5160 taggagaaac gggtagtagg aatttgtaca tttggatcgg aaagtcattc tctttggata 5220 acaactgttc tctagtagat agcaacaaag cggcagacac tgtggagaat gttgattggg 5280

```
tacaaattgg tgaatccatt ttgtgtcaga tggacttgcc aaaagatacc cctataaagg 5340
taataatagc ctaaactttg gaggctctga tactttttac taattgtaaa gtctgcgtgc 5400
tcatctttgt catgtcttat ccaaccaaac tatatttcga agatgaaaat tacaatctca 5460
gcactttcat tactgactac tgaggacggt taggtagaat ccttatgatt tcagcagttg 5520
tatgtattgg tttattctct agtggtttgc atggttccaa cttgttatga tccttttgtt 5580
gtttgtaact gataagttgc ttttctttct tgttaacaga tagttaggga atctgaggat 5640
cagacagagt tgctagcact gctgagcgcg ctataacacc cacccgcaag ctctacacat 5700
ttactctgtt ttttttcac agattccttc aaccgcaaca cttttccatt ttcagacaga 5760
gtattcattc agetcaggtg agaattctct gaaagcagtc tgtaacactt catcttcaca 5820
gttgcatccg aatacaatcg ttagttctgg attatgttta attgctatct gatcatgaat 5880
ttgagttaga ggatggttgg aacaaaaaaa cttagaagct cgaatgaccg gtttttacca 5940
aatteteata gaccatattt gattettttg atttacttet ggtgeaggae tetetgtget 6000
tatggaagtt gatgttgggg gaaacaactc tcttgtacag tggggaaaaa acttcttctt 6060
cttctttcta tcacatgaaa atcctcaagg gccattatta gtatgatcag attataaaat 6120
tgtaaggtta ggggctttat gaggattttg atggacttgt tacaatgttt acatatacac 6180
tcagcagcac aatagatttt tgttaaactt acatgttatt caagtaaaag tactatgtag 6240
atgttgaagt ctaattgaag aattagttaa tgatagtctt aaacacttga ttcacttgtc 6300
atccaatttt ggttttgcgc atagtttctc ttcttttatt tcctctctaa aacacg
                                                                 6356
<210> 2
<211> 3059
<212> DNA
<213> Arabidopsis thaliana
<400> 2
cttcttctcc gctttgtttc cccaatctct ctcttctctc tctctgaaga aaaataaata 60
aaagatetaa etttgaegge tetettaate ttaeteacte eggttttaag teggaataga 120
ctgatgggag cttgatggtt attgttagat cagatgtgga tttaaagcct tcgctgaact 180
aacaagteta tggaagaage aaagaceett gttttacaet gtatgttgtg aggaatttgt 240
ctgattttgg gtgataaagg tgaagtcttt gagtttgtaa ttttgagata agattggatg 300
gtgggaagag aggatgcgat ggggaatgat gaagctcctc ctggttctaa gaaaatgttt 360
tggcggtctg cctcttggtc tgcttcacgg actgcatcac aagttcctga gggtgatgag 420
caaageetga acatteegtg tgctattagt tetgggeega gtegaagatg teeagetget 480
cetttgacac ctegttcaca tcataacage aaggetagag cttgtttgee accattgeag 540
cctcttgcca tttctaggag gagcttagac gagtggccta aggcgggttc ggatgatgtc 600
ggtgagtggc ctcatccacc aacacctagc gggaacaaaa ccggggagag attgaagctc 660
gatttatcat caacgcagca gcgggtaaca gataagagct ctggtctagc taagagggag 720
aagattgctt tctttgataa agaatgttca aaggttgctg atcatatata tgtgggtgga 780
gatgctgtgg cgaaagacaa gagcatactg aaaaacaatg gaatcacgca tatcttgaac 840
tgcgttggtt ttatctgtcc ggaatatttc aagtctgatt tttgttacag atccttgtgg 900
ttacaggata gtccgtcaga ggatataact agtattctgt acgatgtgtt tgactacttt 960
gaagatgtga gggagcaaag tggaaggatc tttgttcatt gttgtcaagg ggtttcacga 1020
tctacctcgt tggtaatagc atatctgatg tggagagaag ggcaaagttt tgatgatgca 1080
tttcagtatg tgaagtctgc tagaggtatt gctgatccta acatgggctt tgcttgccaa 1140
ttgttacaat gccaaaagag ggttcatgcg ttcccgctta gccctacctc cttacttaga 1200
atgtacaaaa tgtctccaca ctctccttat gaccctttgc atcttgttcc aaaactgttg 1260
aatgatecat geeegggtte tetggattea agaggtgeat ttateattea gttacettet 1320
gcaatttaca tttgggttgg taggcagtgt gaaaccatca tggagaaaga tgcaaaagct 1380
gctgtttgtc agattgctcg atatgagaaa gtcgaagcac ctattatggt ggtcagagaa 1440
ggtgatgagc ctgtttatta ctgggatgca ttttgcaagca ttttgcctat gattggggc 1500
tcggtaatta aagttcaacc aggtgacagg aaggtcgacg catataatct ggattttgag 1560
atttttcaga aagccataga gggaggtttt gtgccaactt tagcatcatc caacaacgaa 1620
catgagactc atcttcctgc aagggaaaac agttggagct cactgaaatg taagtttgca 1680
tcaaggtttg acaaaggttt tcggtatgtc tccaaaacgc cactatctag ggtctattca 1740
gattcgatga tgatcgtgca ttcatcaggc tcaccttcct caacaacttc ttcatcatcc 1800
```

-4-

```
actgegtege etecttitet etetecegat tetgtatgtt caacaaatte aggeaatage 1860
ttaaagagtt tototoaato ototggacgt togtoottga gacottotat tocaccatog 1920
ctgacattgc ctaaattttc cagcctatcc ctcctccctt cccaaacttc tcctaaagaa 1980
tctcgtggtg tcaatacttt tcttcaaccg tcaccaaata gaaaggcttc accttctctt 2040
gergagegta gaggeageet gaaaggatet etgaagttge eaggtttgge tgatteeaac 2100
agaggcacac ctgcttttac tttacatccg gatgatagta atgacatagt cttcaatctg 2160
gagggtatta gaaacggcga tctatatcca ccaagtgatt gcaaagggac aagtgtagat 2220
tcagatttgc cagagaagga aattatatcc ttaatcagtt qcagtaaatc tgacagacac 2280
aaategggag gtgatactga tagetetgge cageetttag catgtegttg gecaagtatg 2340
gagatgatta caaaactgag cagagcttac ttagattcag aatctgttat agcaatcccg 2400
ttgccaagtg atgctgtagg agaaacgggt agtaggaatt tgtacatttg gatcggaaag 2460
tcattctctt tggataacaa ctgttctcta gtagatagca acaaagcggc agacactgtg 2520
gagaatgttg attgggtaca aattggtgaa tecattttgt gtcagatgga ettgecaaaa 2580
gataccccta taaagatagt tagggaatct gaggatcaga cagagttgct agcactgctg 2640
agegegetat aacacccace egeaagetet acacatttae tetgtttttt tttcacagat 2700
tectteaace geaacacttt tecattttea gacagagtat teatteaget caggactete 2760
tgtgcttatg gaagttgatg ttgggggaaa caactctctt gtacagtggg gaaaaaactt 2820
cttcttcttc tttctatcac atgaaaatcc tcaagggcca ttattagtat gatcagatta 2880
taaaattgta aggttagggg ctttatgagg attttgatgg acttgttaca atgtttacat 2940
atacactcag cagcacaata gatttttgtt aaacttacat gttattcaag taaaagtact 3000
<210> 3
<211> 784
<212> PRT
<213> Arabidopsis thaliana
Met Val Gly Arg Glu Asp Ala Met Gly Asn Asp Glu Ala Pro Pro Gly
Ser Lys Lys Met Phe Trp Arg Ser Ala Ser Trp Ser Ala Ser Arg Thr
Ala Ser Gln Val Pro Glu Gly Asp Glu Gln Ser Leu Asn Ile Pro Cys
                           40
Ala Ile Ser Ser Gly Pro Ser Arg Arg Cys Pro Ala Ala Pro Leu Thr
Pro Arg Ser His His Asn Ser Lys Ala Arg Ala Cys Leu Pro Pro Leu
                    70 .
Gln Pro Leu Ala Ile Ser Arg Arg Ser Leu Asp Glu Trp Pro Lys Ala
         85
Gly Ser Asp Asp Val Gly Glu Trp Pro His Pro Pro Thr Pro Ser Gly
                              105
Asn Lys Thr Gly Glu Arg Leu Lys Leu Asp Leu Ser Ser Thr Gln Gln
Arg Val Thr Asp Lys Ser Ser Gly Leu Ala Lys Arg Glu Lys Ile Ala
    130
                       135
                               140
```

Phe 145	Phe	Asp	Lys	Glu	Cys 150	Ser	Lys	Val	Ala	Asp 155	His	Ile	Tyr	Val	Gl ₃ 160
Gly	Asp	Ala	Val	Ala 165	Lys	Asp	Lys	Ser	Ile 170	Leu	Lys	Asn	Asn	Gly 175	Ιlϵ
Thr	His	Ile	Leu 180	Asn	Cys	Val	Gly	Phe 185	Ile	Cys	Pro	Glu	Tyr 190	Phe	Lys
Ser	Asp	Phe 195	Cys	Tyr	Arg	Ser	Leu 200	Trp	Leu	Gln	Asp	Ser 205	Pro	Ser	Glu
Asp	Ile 210	Thr	Ser	Ile	Leu	Tyr 215	Asp	Val	Phe	Asp	Tyr 220	Phe	Glu	Asp	Val
Arg 225	Glu	Gln	Ser	Gly	Arg 230	Ile	Phe	Val	His	Cys 235	Cys	Gln	Gly	Val	Ser 240
Arg	Ser	Thr	Ser	Leu 245	Val	Ile	Ala	Tyr	Leu 250	Met	Trp	Arg	Glu	Gly 255	Glr
Ser	Phe	Asp	Asp 260	Ala	Phe	Gln	Tyr	Val 265	Lys	Ser	Ala	Arg	Gly 270	Ile	Ala
Asp	Pro	Asn 275	Met	Gly	Phe	Ala	Cys 280	Gln	Leu	Leu	Gln	Cys 285	Gln	Lys	Arg
Val	His 290	Ala	Phe	Pro	Leu	Ser 2 9 5	Pro	Thr	Ser	Leu	Leu 300	Arg	Met	Tyr	Lys
Met 305	Ser	Pro	His	Ser	Pro 310	Tyr	Asp	Pro	Leu	His 315	Leu	Val	Pro	Lys	Leu 320
Leu	Asn	Asp	Pro	Cys 325	Pro	Gly	Ser	Leu	Asp 330	Ser	Arg	Gly	Ala	Phe 335	Ile
Ile	Gln	Leu	Pro 340	Ser	Ala	Ile	Tyr	Ile 3 4 5	Trp	Val	Gly	Arg	Gln 350	Cys	Glu
Thr	Ile	Met 355	Glu	Lys	Asp	Ala	Lys 360	Ala	Ala	Val	Cys	Gln 365	Ile	Ala	Arg
Tyr	Glu 370	Lys	Val	Glu	Ala	Pro 375	Ile	Met	Val	Val	Arg 380	Glu	Gly	Asp	Glu
Pro 385	Val	Tyr	Tyr	Trp	Asp 390	Ala	Phe	Ala	Ser	Ile 395	Leu	Pro	Met	Ile	Gly 400
Gly	Ser	Val	Ile	Lys 405	Val	Gln	Pro	Gly	Asp 410	Arg	Lys	Val	Asp	Ala 415	Tyr
Asn	Leu	Asp	Phe 420	Glu	Ile	Phe	Gln	Lys 425	Ala	Ile	Glu	Gly	Gly 430	Phe	Val
Pro	Thr	Leu	Ala	Ser	Ser	Asn	Asn	Glu	Hie	Glu	مددال	Wie	Len	Dro	אן ב

-6**-**

			435					440					445			
P	urg	Glu 450	Asn	Ser	Trp	Ser	Ser 455	Leu	Lys	Cys	Lys	Phe 460	Ala	Ser	Arg	Phe
	usp 165	Lys	Gly	Phe	Arg	Tyr 470	Val	Ser	Lys	Thr	Pro 475	Leu	Ser	Arg	Val	Тут 480
2	er	Asp	Ser	Met	Met 485	Ile	Val	His		Ser 490	Gly	Ser	Pro	Ser	Ser 495	Thr
ľ	hr	Ser	Ser	Ser 500	Ser	Thr	Ala		Pro 505		Phe	Leu	Ser	Pro 510	Asp	Ser
V	/al	Cys	Ser 515	Thr	Asn	Ser	Gly	Asn 520	Ser	Leu	Lys	Ser	Phe 525	Ser	Gln	Ser
2	Ser	Gly 530	Arg	Ser	Ser	Leu	Arg 535	Pro	Ser	Ile	Pro	Pro 540	Ser	Leu	Thr	Leu
	Pro 545	Lys	Phe	Ser	Ser	Leu 550	Ser	Leu	Leu	Pro	Ser 555	Gln	Thr	Ser	Pro	Lys 560
G	Slu	Ser	Arg	Gly	Val 565	Asn	Thr	Phe	Leu	Gln 570	Pro	Ser	Pro	Asn	Arg 575	Lys
P	la	Ser	Pro	Ser 580	Leu	Ala	Glu	Arg	Arg 585	Gly	Ser	Leu	Lys	Gly 5 9 0	Ser	Leu
I	ys	Leu	Pro 595	Gly	Leu	Ala	Asp	Ser 600	Asn	Arg	Gly	Thr	Pro 605	Ala	Phe	Thr
Ι	.eu	His 610	Pro	Asp	Asp	Ser	Asn 615	Asp	Ile	Val	Phe	Asn 620	Leu	Glu	Gly	Ile
	arg 525	Asn	Gly	Asp	Leu	Tyr 630	Pro	Pro	Ser	Asp	Cys 635	Lys	Gly	Thr		Val 640
I	/sp	Ser	Asp	Leu	Pro 645	Glu	Lys	Glu	Ile	Ile 650	Ser	Leu	Ile	Ser	Cys 655	Ser
I	Lys	Ser	Asp	Arg 660	His	Lys	Ser	Gly	Gly 665	Asp	Thr	Asp	Ser	Ser 670	Gly	Gln
Ι	Pro	Leu	Ala 675	Cys	Arg	Trp	Pro	Ser 680	Met	Glu	Met	Ile	Thr 685	Lys	Leu	Ser
2	Arg	Ala 690	Tyr	Leu	Asp	Ser	Glu 695	Ser	Val	Ile	Ala	Ile 700	Pro	Leu	Pro	Ser
_	Asp 705	Ala	Val	Gly	Glu	Thr 710	Gly	Ser	Arg	Asn	Leu 715	Tyr	Ile	Trp	Ile	Gly 720
1	Ĺys	Ser	Phe	Ser	Leu 725	Asp	Asn	Asn	Cys	Ser 730	Leu	Val	Asp	Ser	Asn 735	Lys

-1-7-

```
Ala Ala Asp Thr Val Glu Asn Val Asp Trp Val Gln Ile Gly Glu Ser
                                745
Ile Leu Cys Gln Met Asp Leu Pro Lys Asp Thr Pro Ile Lys Ile Val
                            760
Arg Glu Ser Glu Asp Gln Thr Glu Leu Leu Ala Leu Leu Ser Ala Leu
                        775
<210> 4
<211> 10
<212> PRT
<213> Arabidopsis thaliana
<400> 4
Val His Cys Cys Gln Gly Val Ser Arg Ser
<210> 5
<211> 10
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: mammalian
      sequence motif defining the family of PTPs
<400> 5
Ile His Cys Xaa Ala Gly Xaa Xaa Arg Ser
<210> 6
<211> 14
<212> PRT
<213> Arabidopsis thaliana
<400> 6
Thr Ser Ile Leu Tyr Asp Val Phe Asp Tyr Phe Glu Asp Val
  1
                                     10
<210> 7
<211> 12
<212> PRT
<213> Arabidopsis thaliana
<400> 7
Phe Val His Cys Cys Gln Gly Val Ser Arg Ser Thr
```

-8-

```
1
                  5
                                      10
<210> 8
<211> 4
<212> PRT
<213> Arabidopsis thaliana
<400> 8
Phe Val His Cys
 1
<210> 9
<211> 6
<212> PRT
<213> Arabidopsis thaliana
<400> 9
Gln Gly Val Ser Arg Ser
                  5
<210> 10
<211> 5
<212> PRT
<213> Arabidopsis thaliana
<400> 10
Tyr Phe Lys Ser Asp
 1
<210> 11
<211> 23
<212> DNA
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: degenerate
     primer
<400> 11
aayaayggna thacncayat hyt
                                                                    23
<210> 12
<211> 24
<212> DNA
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: degenerate
     primer
<400> 12
```

greanger aanceeatrt tngg	24
10> 13 1> 24 2> DNA _13> Artificial Sequence	
<220> <223> Description of Artificial Sequence: degenerate primer	
<400> 13	
ngtccacatn arrtangcda tnac	24
<210> 14 <211> 21 <212> DNA <213> Artificial Sequence	
<220> <223> Description of Artificial Sequence: primer	
<400> 14 ggtttctaca ggacgtaaca t	21
<210> 15 <211> 196 <212> DNA <213> Sinapis alba	
<400> 15 caactgegtt ggttteatet gteetgaata ttteaagtet gatttttgtt aeeggtegtt gtggttaegt gatagteeat eagaggatat aaetageatt etetaegatg tetttgaeta etttgaagae gttagggage aaagtgggag gatetttgtt eaetgttgte aaggegttte aeggtetaee teettg	120
<210> 16 <211> 65 <212> PRT <213> Sinapis alba	
<pre><400> 16 Asn Cys Val Gly Phe Ile Cys Pro Glu Tyr Phe Lys Ser Asp Phe Cys 1</pre>	
Tyr Arg Ser Leu Trp Leu Arg Asp Ser Pro Ser Glu Asp Ile Thr Ser 20 25 30	~ *
Ile Leu Tyr Asp Val Phe Asp Tyr Phe Glu Asp Val Arg Glu Gln Ser 35 40 45	
Gly Arg Ile Phe Val His Cys Cys Gln Gly Val Ser Arg Ser Thr Ser 50 55 60	
Leu	

. a - 10 -

```
65
<210> 17
<211> 196
<212> DINA
<213> Lycopersicon esculentum
<400> 17
aaactgtgta gggtttagtt gtcctgaata ctttaaggat gaccttgtat acaagacact 60
ttggctgcag gatagcccca ctgaggacat cacaagtatt ctttatgatg tctttgatta 120
ctttgaagat gttcatgaac aaggtgggag tgtctttgta cactgcttcc agggggtgtc 180
ccgatcagcc tccttg
<210> 18
<211> 65
<212> PRT
<213> Lycopersicon esculentum
<400> 18
Asn Cys Val Gly Phe Ser Cys Pro Glu Tyr Phe Lys Asp Asp Leu Val
Tyr Lys Thr Leu Trp Leu Gln Asp Ser Pro Thr Glu Asp Ile Thr Ser
                                 25
Ile Leu Tyr Asp Val Phe Asp Tyr Phe Glu Asp Val His Glu Gln Gly
Gly Ser Val Phe Val His Cys Phe Gln Gly Val Ser Arg Ser Ala Ser
                         55
Leu
 65
<210> 19
<211> 196
<212> DNA
<213> Zea mays
<400> 19
caactgcatg ggcttcgtct gcccgaagta cttcaagtct gaccttgtct accgcaccct 60
ctggctgcag gacagcccca ccgaggacac caccagcatc ctttacgacg tgtttgatta 120
ctttgaggac gtcagggagc agggtggccg cgtgtttgtg cattgctgcc agggggtgtc 180
gcgctccacg cctctg
<210> 20
<211> 65
<212> PRT
<213> Zea mays
<400> 20
Asn Cys Met Gly Phe Val Cys Pro Lys Tyr Phe Lys Ser Asp Leu Val
```

10

- 11 -

```
Tyr Arg Thr Leu Trp Leu Gln Asp Ser Pro Thr Glu Asp Thr Thr Ser
Ile Leu Tyr Asp Val Phe Asp Tyr Phe Glu Asp Val Arg Glu Gln Gly
                            40
Gly Arg Val Phe Val His Cys Cys Gln Gly Val Ser Arg Ser Thr Pro
Leu
65
<210> 21
<211> 2450
<212> DNA
<213> Zea mays
<400> 21
ggcgggtcct ccccgccaa gcccggggag ggctccgcc tcgacctctc ctcgctccgg 60
tegeagggge geaaggacea gategeette ttegacaagg agtgeteeaa ggtegeegae 120
cacgtctacc tcggcggcga cgccgtcgcc aagaaccgcg acatcctcag gaagaacggc 180
atcacccacg tgctcaactg tgtgggcttt gtctgcccgg agtacttcaa gtcggaccta 240
gtotacogca coctotggot goaggacago cocacogagg acatoacoag catootgtac 300
gacgtgtttg actacttcga ggacgtcagg gagcagggcg gccgcgtgct tgtgcattgc 360
tgecaggggg tgtcacgctc cacgtcgctg gtcatcgcct acttgatgtg gagggaaggc 420
cagagetteg atgacgeett ceagtttgte aaggetgeee gggggatege aaatecaaac 480
atgggetttg catgecaget tetecagtge cagaagegtg tgcatgegat teegetgtca 540
ccaaattcag tgctcaggat gtaccgcatg gcgcctcact cccagtatgc ccctctgcat 600
ttggtgccca aaatgctcaa tgacccatcc ccagccaccc ttgactctag aggcgcgttc 660
attgtgcatg ttctctcgtc gctctatgtc tgggttggaa tgaagtgtga tccggtaatg 720
gaaaaggatg caaaggctgc tgcgtttcag gtagtgaggt atgagaaggt gcaggggcac 780
atcaaggttg tgagagaagg tctggagccg caggagttct gggatgcatt ttcaagtatg 840
ccacctaatt cagatagtaa tacaaagatt agcaaggacc agatcgattc agcatccaag 900
agtgacccag gaagccggaa aaatgagtcc tatgacgctg attttgagct tgtctacaaa 960
gcaatcactg ggggagtggt ccctgcattt tcaacttctg gggctggtga tgagacccat 1020
cttccagcta gagaaagtag ctggagttta ctgaggcaca agtttatctc caggtcgcta 1080
gctcgtgttt attcagattc tgctctaatg aaggattttg atccacgggt acaacacctg 1140
getgetgagg catcaacete aceteette etttetecaa geteettate ateggattea 1200
agtgtcaget egaagtatag tteagaetea eeeteettat eacetacaae tggeteteca 1260
ccatcatttg gcctctcgcc tgcttcatct aatctgacac atgctttggt gccatcatcc 1320
aggteteece ttteteaate atetaatgaa ggagetteaa ageettetgg catggaatea 1380
atacactete ettecaagae etettetata geagaaagga gaggaggett cacactteta 1440''
ccaccaaggg cgccatctag tattcgcagg accgaggatg cctcagataa tagtacaaat 1560
ggggttaaac agctgactag tgagttttgc tcagaaaaat gcactggtaa tagtttgagc 1620
tegeattetg aaactagatt aattgagegt actgacagta actcagaagt ctgcagtaat 1680
gcacaacttg tagtctacca gtggcccagc atgggaaagc taactacatt tgcacgcaag 1740
gatettgace egaagteggt tttaattttt gttaettega atgecateag gagaggagaa 1800
gcagttaaaa tggtgtatgt atgggtagga ggcgaaaatg agagcagcaa gagtgttgac 1860
tccgtcgatt ggcaacaggt cactagtgat tttcttcatc taaagggcct cagcaatgtt 1920
cttcctgtca aggttttcaa ggagcatgaa gctgagaatc ttttggaact actgaatgtt 1980
agttaacatt aggcagtagc tatcaggata attgtagttg ctaaacaaac tcaacgaagg 2040
```

The Court

- - - VI

```
catgecetee ageateagte ggtacegatg attgtcageg aggtataaag ccaeagecat 2100
tcccttgaac ataataagct acaaacagat tccgttctgc aactgcgcct catgatctat 2160
attitgtcca gatggcagga ggctgccatg ggcgttgtat cggttgcgaa ttagcactcg 2220
tggtgtagga gcaatcggcc gattcggtgt atattatccg ctcccctgta atgtaagctc 2280
agatactggg agctggtgtg tcgacagtta ctttttagcc taaacattct tgtacatctt 2340
tqaaaqqaac agagttgtaa teettttgac tatgtaaatg geteeattgg teataaette 2400
2450
<210> 22
<211> 661
<212> PRT
<213> Zea mays
<400> 22
Gly Gly Ser Ser Pro Ala Lys Pro Gly Glu Gly Leu Arg Leu Asp Leu
Ser Ser Leu Arg Ser Gln Gly Arg Lys Asp Gln Ile Ala Phe Phe Asp
Lys Glu Cys Ser Lys Val Ala Asp His Val Tyr Leu Gly Gly Asp Ala
Val Ala Lys Asn Arg Asp Ile Leu Arg Lys Asn Gly Ile Thr His Val
Leu Asn Cys Val Gly Phe Val Cys Pro Glu Tyr Phe Lys Ser Asp Leu
Val Tyr Arg Thr Leu Trp Leu Gln Asp Ser Pro Thr Glu Asp Ile Thr
Ser Ile Leu Tyr Asp Val Phe Asp Tyr Phe Glu Asp Val Arg Glu Gln
                               105
Gly Gly Arg Val Leu Val His Cys Cys Gln Gly Val Ser Arg Ser Thr
Ser Leu Val Ile Ala Tyr Leu Met Trp Arg Glu Gly Gln Ser Phe Asp
                       135
Asp Ala Phe Gln Phe Val Lys Ala Ala Arg Gly Ile Ala Asn Pro Asn
145
Met Gly Phe Ala Cys Gln Leu Leu Gln Cys Gln Lys Arg Val His Ala
                                   170
Ile Pro Leu Ser Pro Asn Ser Val Leu Arg Met Tyr Arg Met Ala Pro
                               185
            180
His Ser Gln Tyr Ala Pro Leu His Leu Val Pro Lys Met Leu Asn Asp
Pro Ser Pro Ala Thr Leu Asp Ser Arg Gly Ala Phe Ile Val His Val
```

225	ser	Ser	Deu	TYT	230	пр	Val	GIY	Mec	235	cys	rsp	PIO	vai	240	
Glu	Lys	Asp	Ala	Lys 245	Ala	Ala	Ala	Phe	Gln 250	Val	Val	Arg	Tyr	Glu 255	Lys	
Val	Gln	Gly	His 260	Ile	Lys	Val	Val	Arg 265	Glu	Gly	Leu	Glu	Pro 270	Gln	Glu	
Phe	Trp	Asp 275	Ala	Phe	Ser	Ser	Met 280	Pro	Pro	Asn	Ser	Asp 285	Ser	Asn	Thr	
Lys	Ile 290	Ser	Lys	Asp	Gln	Ile 295	Asp	Ser	Ala	Ser	Lys 300	Ser	Asp	Pro	Ğly	
Ser 305	Arg	Lys	Asn	Glu	Ser 310	Tyr	Asp	Ala	Asp	Phe 315	Glu	Leu	Val	Tyr	Lys 320	
Ala	Ile	Thr	Gly	Gly 325	Val	Val	Pro	Ala	Phe 330	Ser	Thr	Ser	Gly	Ala 335	Gly	
Asp	Glu	Thr	His 3 4 0	Leu	Pro	Ala	Arg	Glu 3 4 5	Ser	Ser	Trp	Ser	Leu 350	Leu	Arg	
His	Lys	Phe 355	Ile	Ser	Arg	Ser	Leu 360	Ala	Arg	Val	Tyr	Ser 365	Asp	Ser	Ala	
Leu	Met 370	Lys	Asp	Phe	Asp	Pro 375	Arg	Val	Gln	His	Leu 380	Ala	Ala	Glu	Ala	
Ser 385	Thr	Ser	Pro	Pro	Phe 390	Leu	Ser	Pro	Ser	Ser 395	Leu	Ser	Ser	Asp	Ser 400	
Ser	Val	Ser	Ser	Lys 405	Tyr	Ser	Ser	Asp	Ser 410	Pro	Ser	Leu	Ser	Pro 415	Thr	
Thr	Gly	Ser	Pro 420	Pro	Ser	Phe	Gly	Leu 425	Ser	Pro	Ala	Ser	Ser 430	Asn	Leu	
Thr	His	Ala 435	Leu	Val	Pro	Ser	Ser 440	_	Ser	Pro	Leu	Ser 445	Gln	Ser	Ser	
Asn	Glu 450	Gly	Ala	Ser	Lys	Pro 455	Ser	Gly	Met	Glu	Ser 460	Ile	His	Ser	Pro	
Ser 465	Lys	Thr	Ser	Ser	Ile 470	Ala	Glu	Arg	Arg	Gly 475	Gly	Phe	Thr	Leu	Leu 480	
Lys	Leu	Pro	Ser	Leu 485	Gln	Lys	Asp	Leu	Val 490	Leu	Pro	Pro	Arg	Val 495	Pro	
Ser	Ile	Val	Leu 500		Pro	Arg	Ala	Pro		Ser	Ile	Arg	Arg		Glu	

1.77. .22.

-8t - -14 -

```
Asp Ala Ser Asp Asn Ser Thr Asn Gly Val Lys Gln Leu Thr Ser Glu
        515
                            520
                                                 525
Phe Cys Ser Glu Lys Cys Thr Gly Asn Ser Leu Ser Ser His Ser Glu
                        535
Thr Arg Leu Ile Glu Arg Thr Asp Ser Asn Ser Glu Val Cys Ser Asn
                    550
                                        555
Ala Gln Leu Val Val Tyr Gln Trp Pro Ser Met Gly Lys Leu Thr Thr
                565
                                    570 .
Phe Ala Arg Lys Asp Leu Asp Pro Lys Ser Val Leu Ile Phe Val Thr
                                585
Ser Asn Ala Ile Arg Arg Gly Glu Ala Val Lys Met Val Tyr Val Trp
        595
                            600
                                                 605
Val Gly Glu Asn Glu Ser Ser Lys Ser Val Asp Ser Val Asp Trp
                        615
Gln Gln Val Thr Ser Asp Phe Leu His Leu Lys Gly Leu Ser Asn Val
                                        635
Leu Pro Val Lys Val Phe Lys Glu His Glu Ala Glu Asn Leu Leu Glu
                645
                                    650
Leu Leu Asn Val Ser
            660
<210> 23
<211> 522
<212> DNA
<213> Lycopersicon esculentum
<400> 23
aactgtgtgg ggtttgtatg cccagagtat ttcaagtctg atttcgtata ccggactttg 60
tggttgcagg atagcccatc agaagatatt actagtattc tctatgatgt ttttgactac 120
tttgaagatg tcagggagca acatgggaag gtttttgttc attgctgcca aggggtctct 180
eggteaacct egttggttat tgettategt atgtggagag aaggacaaag ttttgatgat 240
geetttgagt atgtaaagge ageaaggggt attgeggate caaatatggg ttttgettgt 300
cagttattac aatgccaaaa aagggttcat gcttctcctt tgagcccaag ttcattatta 360
aggatgtaca gagttgcacc tcattcacca tacgatcctt tgcatctcgt cccaaaaatg 420
ttaaatgatc cctcaccggc agcattagat tctagaggtg catttattat acacatacct 480
tcatcggtat atgtatggat tggtaagaaa tgtgaagcaa tc
<210> 24
<211> 174
<212> PRT
<213> Lycopersicon esculentum
<400> 24
Asn Cys Val Gly Phe Val Cys Pro Glu Tyr Phe Lys Ser Asp Phe Val
                                     10
```

Tyr Ai	g Thi	20 20	ŢŢ	Leu	Gln	Asp	Ser 25	Pro	Ser	Glu	Asp	Ile 30	Thr	Ser	
Ile Le	eu Tyr 35	_	Val	Phe	Asp	Tyr 40	Phe	Glu	Asp	Val	Arg 45	Glu	Gln	His	
Gly Ly	vs Val	. Phe	Val	His	Cys 55	Cys	Gln	Gly	Val	Ser 60	Arg	Ser	Thr	Ser	
Leu Va 65	al Il∈	Ala	Tyr	Arg 70	Met	Trp	Arg	Glu	Gly 75	Gln	Ser	Phe	Asp	Asp 80	
Ala Ph	ne Glu	Tyr	Val 85	Lys	Ala	Ala	Arg	Gly 90	Ile	Ala	Asp	Pro	Asn 95	Met	
Gly Ph	ne Ala	Cys 100	Gln	Leu	Leu	Gln	Cys 105	Gln	Lys	Arg	Val	His 110	Ala	Ser	
Pro Le	u Ser 115		Ser	Ser	Leu	Leu 120	Arg	Met	Tyr	Arg	Val 125	Ala	Pro	His	
Ser Pr		: Asp	Pro	Leu	His 135	Leu	Val	Pro	Lys	Met 140	Leu	Asn	Asp	Pro	٠
Ser Pr 145	o Ala	Ala	Leu	Asp 150	Ser	Arg	Gly	Ala	Phe 155	Ile	Ile	His	Ile	Pro 160	
Ser Se	er Val	. Tyr	Val 165	Trp	Ile	Gly	Lys	Lys 170	Cys	Glu	Ala	Ile			
<210> <211> <212> <213>	21 DINA	icia	l Sec	quenc	ce										
<220> <223>	Descr	riptio	on of	E Art	ific	cial	Sea	ience	e: de	ecrene	erate	<u> -</u>			
	prime						•								
<400> gangay		crtc	yttyt	c c											21
<210> <211> <212> <213>	24 DINA	icia	l Sec	quenc	ce										
<220> <223>	Descr prime		on of	E Art	ific	cial	Seqi	ıence	e: d	egene	erate	9			
<400> ytcnck		ggna	rrtgi	ng ty	ytc										24

WORLD INTELLECTUAL PROPERTY ORGANIZATION International Bureau



INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

(51) International Patent Classification 7: WO 00/06706 (11) International Publication Number: **A3** C12N 9/16 (43) International Publication Date: 10 February 2000 (10.02.00)

PCT/EP99/05413 (21) International Application Number:

28 July 1999 (28.07.99) (22) International Filing Date:

(30) Priority Data:

9816639.0

30 July 1998 (30.07.98)

GB

(71) Applicant (for all designated States except AT US): NOVAR-TIS AG [CH/CH]; Schwarzwaldallee 215, CH-4058 Basel (CH).

(71) Applicant (for AT only): NOVARTIS-ERFINDUNGEN VER-WALTUNGSGESELLSCHAFT MBH [AT/AT]; Brunner | Published Strasse 59, A-1230 Vienna (AT).

(72) Inventors; and

- (75) Inventors/Applicants (for US only): REVENKOVA, Ekaterina [RU/CH]; Peter Rot-Strasse 76, CH-4058 Basel (CH). PASZOWSKI, Jerzy [PL/CH]; Blauenweg 10, CH-4224 Nenzlingen (CH).
- (74) Agent: BECKER, Konrad; Novartis AG, Corporate Intellectual Property, Patent & Trademark Dept., CH-4002 Basel (CH).

(81) Designated States: AE, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CU, CZ, DE, DK, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, UA, UG, US, UZ, VN, YU, ZA, ZW, ARIPO patent (GH, GM, KE, LS, MW, SD, SL, SZ, UG, ZW), Eurasian patent (AM, AZ, BY, KG, KZ, MD, RU, TJ, TM), European patent (AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE), OAPI patent (BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG).

With international search report.

Before the expiration of the time limit for amending the claims and to be republished in the event of the receipt of amendments.

(88) Date of publication of the international search report: 4 May 2000 (04.05.00)

(54) Title: MAP KINASE PHOSPHATASE MUTANT

(57) Abstract

The present invention relates to DNA encoding proteins contributing to the regulation of a plant's response to DNA damage. DNA according to the invention comprises an open reading frame encoding a protein characterized by a stretch of amino acids or component amino acid sequence having 40% or more identity with an aligned component sequence of SEQ ID NO:3. Preferably the DNA encodes a MAP kinase phosphatase.

FOR THE PURPOSES OF INFORMATION ONLY

Codes used to identify States party to the PCT on the front pages of pamphlets publishing international applications under the PCT.

AL	Albania	ES	Spain	LS	Lesotho	SI	Slovenia
AM	Armenia	FI	Finland	LT	Lithuania	SK	Slovakia
AT	Austria	FR	France	LU	Luxembourg	SN	Senegal
AU	Australia	GA	Gabon	LV	Latvia	SZ	Swaziland
ΑZ	Azerbaijan	GB	United Kingdom	MC	Моласо	TD	Chad
BA	Bosnia and Herregovina	GE	Georgia	MD	Republic of Moldova	TG	Togo
BB	Barbados	GII	Ghana	MG	Madagascar	Ŧj	Tajikistan
BE	Belgium	GN	Guinea	MK	The former Yugoslav	TM	Turkmenisten
BF	Burkina Faso	GR	Greece		Republic of Macedonia	TR	Turkey
BG	Bulgaria	ΠÜ	Hungary	ML	Mali	TT	Trinidad and Tohage
BJ	Benin	IE	Ireland	MN	Mongolia	UA	Ukraine
BR	Brazil	IL	Israel	MR	Mauritania	UG	Uganda
BY	Belarus	IS	Iceland	MW	Malawi	US	United States of America
CA	Canada	IT	Italy	MX	Mexico	UZ	Uzbekistan
CF	Central African Republic	JР	Japan	NE	Niger	VN	Vict Nam
CG	Congo	KE	Kenya	NL	Netherlands	YU	Yugoslavia
CH	Switzerland	KG	Kyrgyzstan	NO	Norway	zw	Zimbabwe
CI	Côte d'Ivone	KP	Democratic People's	NZ	New Zealand	2	Zimbabwe
CM	Cameroon		Republic of Korea	PL	Poland		
CN	China	KR	Republic of Korea	PT	Portugal		
CU	Cuba	KZ	Kazakstan	RO	Romania		
CZ	Czech Republic	LC	Saint Lucia	RU	Russian Federation		
DE	Germany	1.1	Liechtenstein	SD	Sudan		
ÐK	Denmark	LK	Sri Lanka	SE	Sweden		- •
EE	Estonia	LR	Liberia	SG	Singapore		

INTERNATIONAL SEARCH REPORT

onel Application No

PCT/EP 99/05413 A. CLASSIFICATION OF SUBJECT MATTER IPC 7 C12N9/16 According to international Patent Classification (IPC) or to both national classification and IPC **B. FIELDS SEARCHED** Minimum documentation searched (classification system followed by classification symbols) IPC 7 C12N Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched Electronic data base consulted during the international search (name of data base and, where practical, search terms used) C. DOCUMENTS CONSIDERED TO BE RELEVANT Category * Citation of document, with indication, where appropriate, of the relevant passages Relevant to claim No. X WO 94 23039 A (CANCER RES INST ROYAL 1-12 ;MARSHALL CHRISTOPHER JOHN (GB); ASHWORTH AL) 13 October 1994 (1994-10-13) figure 7D X & DATABASE DGENE 'Online! 1-12 DERWENT Accession no. R63602, 9 May 1995 (1995-05-09) abstract X DATABASE SWISSPROT 'Online! 1-12 Accession no. P28562, 1 December 1992 (1992-12-01) KEYES S.M. AND EMSLIE E.A.: "DUAL SPECIFICITY PROTEIN PHOSPHATASE 1 XP002131991 abstract -/--X Further documents are listed in the continuation of box C. Patent family members are listed in annex.

* Special categories of	cited documents	

- "A" document defining the general state of the art which is not considered to be of particular relevance
- "E" earlier document but published on or after the International filing date
- "L" document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another claston or other special reason (as specified) e of another
- "O" document referring to an oral disclosure, use, exhibition or other means
- document published prior to the international filing date but later than the priority date claimed
- "I" later document published after the international filing date or priority date and not in conflict with the application but clear to understand the principle or theory underlying the invention
- "X" document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone

. ..

"Y" document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such documents, such combination being obvious to a person skilled in the art.

Date of mailing of the international search report

"&" document member of the same patent family

Date of the actual completion of the international search

14/03/2000

Name and mailing address of the ISA

1 March 2000

European Patent Office, P.B. 5818 Patentiaan 2 NL - 2280 HV Rijswijk Tel. (+91-70) 940-2040, Tx. 91 651 epo ni, Fax: (+91-70) 940-3016

Authorized officer

Chakravarty, A

Form PCT/ISA/210 (second sheet) (July 1992)

1

INTERNATIONAL SEARCH REPORT

Inte. onal Application No
PCT/EP 99/05413

		PCT/EP 9	9/05413
	ation) DOCUMENTS CONSIDERED T BE RELEVANT		
Category *	Citation of document, with indication, where appropriate, of the relevant passages		Relevant to claim No.
(DATABASE SWISSPROT 'Online! Accession no. P28563, 1 December 1992 (1992-12-01) CHARLES C.H. ET AL.: "DUAL SPECIFICITY PROTEIN PHOSPHATASE 1 (EC 3.1.3.48) (EC 3.1.3.16)" XP002131992 cited in the application abstract		1-12
, X	DATABASE EMBL 'Online! EBI accession no. AL085836, 28 June 1999 (1999-06-28) SALANOUBAT M. ET AL.: "T7 end of BAC F1115 of IGF library from strain Columbia of Arabidopsis thaliana" XP002131993 abstract		1-12
			

INTERNATIONAL SEARCH REPORT

information on patent family members

PCT/EP 99/05413

Patent document cited in search report		Publication date		Patent family member(s)	Publication date		
WO 9423039	Α	13-10-1994	AU	677834 B	08-05-1997		
			AU	6382394 A	24-10-1994		
			CA	2157774 A	13-10-1994		
			ΕP	0703984 A	03-04-1996		
			JP	9501302 T	10-02-1997		
			US	5958721 A	28-09-1999		
			AU	696939 B	24-09-1998		
			AU	1586195 A	29-08-1995		
			CA	2182967 A	17-08-1995		
			EP	0742827 A	20-11-1996		
			WO	9521923 A	17-08-1995		
			JP	9508795 T	09-09-1997		

were on the second of the second

•

.

,

. .

- · · •